Original Article
Effect of the long non-coding RNA AC007392.4 on growth, invasion and migration for tongue squamous cell carcinoma

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Abstract: Objectives: Long non-coding RNAs (LncRNA) have been reported to affect the development of many types of tumors. However, the research for relationship between LncRNA AC007392.4 and tongue squamous cell carcinoma cells (TSCC) has not been revealed. In this study, we aim to explore the contribution of LncRNA AC007392.4 to growth, invasion and migration in TSCC cells. Methods: To detect LncRNA AC007392.4 expression with QRT-PCR in 20 patients in TSCC tissue and adjacent normal tissue. Then construct pcDNA3.1 (+) expression vector of LncRNA AC007392.4 and transfected into the Cal-27 cell line. By using MTT, transwell migration/invasion assay and apoptosis analysis, we tested the role of LncRNA AC007392.4 on tongue cancer cell growth, migration and invasion. Results: The results indicated that LncRNA AC007392.4 was lower expression in TSCCs tissue than adjacent normal tissue (P<0.05). And overexpression of LncRNA AC007392.4 in Cal-27 cells can promote cell growth (P<0.05) and decrease the cell apoptosis rate (P<0.05), but no effect on cell migration and invasion (P>0.05). Conclusion: We suggest that LncRNA AC007392.4 can influence on TSCC cells growth, but not on invasion and migration, may be an important and useful therapeutic biomarker for TSCC patients in future but need more study.

Keywords: LncRNA AC007392.4, growth, invasion, migration, apoptosis

Introduction

Tongue squamous cell carcinoma (TSCC) is the most common oral squamous cell carcinoma. Its morbidity presents a gradual upward trend in recent years and its five-year survival rate is maintained at about 50% [1]. However, the five-year survival rate of advanced cases is only 27%. At present, its main treatment method is a multidisciplinary integrated sequence therapy focusing on surgical treatment, but surgical treatment often leads to patients’ dysfunction of language, eating and breath etc. and facial deformity with a high recurrence rate and easy metastasis. Invasion and metastasis are important biological properties of tongue squamous carcinoma. The mechanism of tumor infiltration and lymphatic metastasis is a complex process consisting of multiple steps and its concrete mechanism is not clear yet. To seek for a method that not only can cure tumors effectively, but also maintain normal physiological functions of facial appearance, chew and language etc. is a difficult problem that needs to be solved urgently now. At the same time, in practical work, clinicians hope to find out biological indicators which are closely related to occurrence and development of tongue squamous carcinoma and can conduct accurate and objective evaluation on prognosis of tongue squamous cell carcinoma patients to enhance the survival rate and living quality of patients with tongue squamous carcinoma.

According to lengths, non-coding RNA can be divided into two categories: non-coding short-fragment RNA (including shRNA, iRNA and piRNA) and non-coding long-fragment RNA (including Long non-coding RNA and IncRNA). The present research mainly pays attention to non-coding short-fragment RNA such as microRNA and shRNA and research on microR-
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NA in genomes of multiple animals and plants has been reported. However, genomes of mammals can code a great quantity of non-coding long-fragment RNA (LncRNA) which is endogenous RNA which lacks a specific complete open reading frame and has no protein-coding function with a length of more than 200 nucleotides. Originally, LncRNA was considered as a by-product of transcription of RNA polymerase II that had no biological function [2, 3] and was the “noise” of genome transcription. However, research in recent years showed that LncRNA took part in multiple important cell regulation processes. For example, LncRNA participates in x-chromosome silence, genomic imprinting, chromatin modification, transcriptional activation, transcriptional interference and intranuclear transport etc. In addition, in 2011, ‘Cancer Research’ reported that LncRNA was in favor of early diagnosis, prognosis and molecular targeting treatment of tumors with good value of clinical application [4]. Now it is regarded that functions of non-coding long-fragment RNA are as follows [5, 6]: (1) To interfere expression of downstream genes by transcription in the upstream promoter region of protein coding genes; (2) To form complementary double chains with transcript of protein coding genes to interfere shearing of mRNA leading to different shear forms. (3) To affect expression of downstream genes by inhibiting RNA polymerase II or mediate chromatin remodeling and histone modification. (4) To form complementary double chains with transcript of protein coding genes, further generate endogenous shRNA under the action of Dicer for gene expression level regulation. (5) To be a compound of nucleic acid protein formed by structural constituent and proteins. (6) By bonding to special proteins, the LncRNA transcript can regulate activity of corresponding proteins; (7) By bonding to special proteins, to change cytoplasm localization of proteins; (8) To be precursor molecule transcription of micromolecule RNA etc. Research in recent years showed that LncRNA played an important regulating role in occurrence and development of multiple tumors such as liver cancer, lung cancer and breast cancer [7-9]. In the field of tongue squamous carcinoma research, some researchers found low expression of LncMEG3 in tongue squamous carcinoma with bad clinical prognosis, and it could inhibit the cell proliferation and cycle which result in cell apoptosis by overexpression of LncMEG3 in vitro [10]. In addition, by studying UCA1, Fang found high expression of UCA1 in tongue squamous carcinoma and discovered that high expression of UCA1 in cancer indicated metastasis but has nothing to do with hyperplasia [11]. All of these studies indicated that LncRNA might play a closely regulating role in occurrence and development of tongue cancer. Therefore, in this experiment, LncRNA AC007392.4 expression testing was firstly carried out on tongue cancer tissue and corresponding para-carcinoma tissue (20 cases) by QRT-PCR technology and found that there were expression differences in these two types of tissue. Thus, it was determined that AC007392.4 was treated as LncRNA and therapeutic targets of main function research. Moreover, by up-regulating its expression in cell strains of tongue squamous carcinoma, its effects on cell strains of tongue squamous carcinoma were observed.

Materials and methods

Human tissue specimens

Paired primary TSCC samples and adjacent histological normal tissues of the tongue were obtained from 20 patients who were admitted to the Department of Oral and Maxillofacial Surgery of Guangzhou Medical University of Stomatology between May 2012 and August 2013. None of the patients received radiotherapy or chemotherapy or any other treatment prior to surgery.

Quantitative reverse transcription polymerase chain reaction (QRT-PCR)

Total mRNA samples of tongue cancer tissue and adjacent histological normal tissues were prepared with Trizol reagent (Invitrogen, Grand Island, NY 14072, USA) according to the manufacturer’s instructions. First-strand cDNA was synthesized using the GeneAmp® PCR System 9700 (Life Technologies Corporation, USA) with random hexamer primers. The resulting cDNA was amplified by using the SYBR-Green Master PCR Mix (Applied Biosystem, Grand Island, NY 14072, USA) in triplicates. The primer set for amplifying LncRNAs AC007392.4 were in Table 1.

Construction of expression vectors

LncRNA AC007392.4 were amplified by PCR according to its sequence information and
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then inserted into the pcDNA3.1 expression vector (Invitrogen, kpnI and EcoRI as restriction enzyme sites) at a position downstream of the CMV promoter. The cDNA of 293 cell line is used for the template of PCR amplification. The primers used to amplify this fragment are as follows: AC007392.4-F: 5’CGGGGTACCCCCAAGAAGGGTGGTCTTCCAG3’; AC007392.4-R: 5’CCGGAATTCGAAGACAATGACTTTGTGTGTTGC3’. The product size is 765 bps. Then, they were joined together to construct vectors expressing LncRNA AC007392.4.

Cell cultures

TSCC cell lines, Cal-27 and Scc-9 cell lines were obtained from the Department of Oral and Maxillofacial Surgery, Sun Yat-sen Memorial Hospital, Sun Yat-sen University, Guangzhou, China. 293 cell line is purchased from ATCC. The cells were cultured on with 10% fetal bovine serum (FBS, Gibco), incubated at 37°C and supplemented with 5% CO2 in a humidified chamber.

Transient transfection of the Cal-27 cell line

Cal-27 cells were seed onto 24-well plates the day before transfection to ensure 90% confluence at the time of transfection. Transient transfection with 4 μg of pcDNA3.1 or pcDNA3.1-LncRNA AC007392.4 using TurboFect (Life Technologies Corporation, USA) according to the manufacturer’s instructions. Moreover, cells transfection with a negative control pcDNA3.1 named “NC” group and cells left untreated as a control named “Blank” group. The mRNA expression of LncRNA AC007392.4 after transfection was detected by QRT-PCR. The cultures were harvested according to the following experiment design.

Cell proliferation assay

The effects of LncRNA AC007392.4 overexpression on Cal-27 cells proliferation were assessed by using the Cell Proliferation Reagent Kit I (MTT) (Roche, USA). Briefly, the cells were seeded into 96-well plates (5000 cells/ml, 200 µL media per well). After transfection with pcDNA3.1, pcDNA3.1-AC007392.4 was added to each well and documented every 24 h following the manufacturer’s protocol. An MTT solution (5 mg/mL, 20 µL per well) was then added to the cells, and the cells were incubated for 4 h at 37°C. DMSO (150 µL per well) was then added, and the plates were read on an automated microplate spectrophotometer (Bio-Rad, CA, USA) at 492 nm.

Cell migration and invasion assays

To explore the role of LncRNA AC007392.4 on Cal-27 cells migration and invasion, we performed a transwell assay in a 24-well culture plate. For the migration assay, at 24 h after transfection, 1×104 cells in serum-free media were seeded on a fibronectin-coated polycarbonate membrane insert in a transwell apparatus (Costar, Cambridge, MA, USA) and cultured in 1640 media. FBS was added to the lower chamber. For the invasion assays, 1×105 cells in serum-free media were seeded on an insert coated with Matrigel (BD, USA). 1640 Media containing 5% PBS were added into the lower chamber. After 24 h incubation, the cells remaining on the upper membrane were removed with cotton wool, whereas the cells that had migrated and invaded through the membrane were stained with 95% alcohol and 0.1% crystal violet, then imaged and counted using an inverted microscope.

Cell apoptosis analysis

Cal-27 cells transiently transfected with pcDNA3.1 or pcDNA3.1-LncRNA AC007392.4 were harvested at 48 h after transfection. For cell apoptosis analysis, the fixed cells were washed and stained with propidium iodide (PI) and Annexin V-FITC, the cells were analyzed through flow cytometry using the Guava EasyCyte MiniSystem (Keygen, Nanjing, China). All the groups were performed in triplet and statistically analyzed.

Statistical analysis

All experiments were performed three times in triplicates. The data were analyzed with
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Results

LncRNA AC007392.4 expression was reduced in TSCC versus adjacent histological normal tissues

QRT-PCR analysis was performed on twenty paired samples of TSCC versus matched adjacent histological normal tissues to analyze differential expression of LncRNA AC007392.4. The average expression level of LncRNA AC007392.4 was significantly reduced in TSCC versus adjacent histological normal tissues (Figure 1A, *P<0.05). Furthermore, we tested the differential expression of LncRNA AC007392.4 between Cal-27 and Scc-9 cell lines and found LncRNA AC007392.4 mRNA expression in the Cal-27 cell line was significantly decreased compared with Scc-9 cell line (Figure 1B, *P<0.05).

Effect of LncRNA AC007392.4 overexpression on Cal-27 cells proliferation

We detected whether LncRNA AC007392.4 overexpression affected the proliferation ability of tongue cancer cells by MTT assay. Cal-27 cells transfected with pcDNA3.1 or pcDNA3.1-LncRNA AC007392.4 were harvested at 24 h,
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48 h and 72 h after transfection. LncRNA AC007392.4 overexpression showed higher growth rate compared with the NC and Blank groups at 48 h and 72 h (Figure 3, *P<0.05). These results revealed that LncRNA AC007392.4 overexpression in tongue cancer cells did have a positive effect on cell proliferation.

Effects of LncRNA AC007392.4 overexpression on Cal-27 cells migration and invasion

To explore whether tongue cancer cell migration and invasion were affected by LncRNA AC007392.4 overexpression, a transwell assay was performed. Figure 4A showed the pictures of crystal violet staining cells in different groups. Moreover, there were no significant differences in the LncRNA AC007392.4, NC and Blank groups (Figure 4B, *P>0.05). In this part, we could conclude that possibly LncRNA AC007392.4 do not play any role in the migration and invasion of tongue cancer cell.

Effect of LncRNA AC007392.4 overexpression on Cal-27 cells apoptosis

To investigate the effect of LncRNA AC007392.4 overexpression on Cal-27 cell apoptosis, cells from LncRNA AC007392.4, NC and Blank groups were subjected to Annexin V/PI staining. The apoptosis rate in the LncRNA AC007392.4 group was significantly lower compared with the NC and Blank groups (Figure 5, *P<0.05).

Discussion

The transcribed ucRNAs (T-UCRs) were first described several years ago and they are relevant to some diseases, such as tumor, cardiovascular diseases, etc. Moreover, it has been found that T-UCRs could be regulated by microRNAs [12]. Some reports have been demonstrated that some LncRNAs play important roles in the in TSCC biology [10]. LncRNAs cannot code proteins, but it plays an important role in high order chromosome dynamics, telomere biology and subcellular structure organization [13]. Evidences showed that LncRNAs could regulate gene expression by chromosome modifications and regulation of gene transcription levels and post-transcriptional level through epigenetics to influence biological behaviour of tumor cells [14-16]. Therefore, this experiment intended to explore and study biological behavior, including proliferation, migration, infiltration and apoptosis, of tongue squamous cell carcinoma by expression of LncRNAs AC007392.4.

To expand the functional characterization of LncRNA AC007392.4, we identified LncRNA AC007392.4 was lower expression in TSCCs tissue than adjacent normal tissue via QRT-PCR. Thus, we hypothesized that LncRNA
AC007392.4 could act as an oncogene in TSCC and study the function of LncRNA AC007392.4 by the overexpression. First of all, we tested the differential expression of LncRNA AC007392.4 between Cal-27 and Scc-9 cell lines and found LncRNA AC007392.4 mRNA expression in the Cal-27 cell line was significantly decreased compared with Scc-9 cell line. So we chose Cal-27 cell line for the transfection on the expression of LncRNA AC007392.4 in order to examine and understand the role of LncRNA AC007392.4 directly.

**Figure 4.** Effects of LncRNA AC007392.4 overexpression on Cal-27 cells migration and invasion were performed by transwell assay. A. The pictures of crystal violet staining cells in different groups. B. There were no significant differences in the LncRNA AC007392.4, NC and Blank groups (P>0.05).

**Figure 5.** Effect of LncRNA AC007392.4 overexpression on Cal-27 cells apoptosis was performed by fluorescence-activated cell sorting (FACS) analysis. The apoptosis rate in the LncRNA AC007392.4 group was significantly lower compared with the NC and Blank groups (*P<0.05).
Our results showed that overexpression of LncRNA AC007392.4 in Cal-27 cells can promote cell growth (through MTT analysis) and decrease the cell apoptosis rate, but no effect on cell migration and invasion. Above all, we could conclude that LncRNA AC007392.4 has influence on the TSCC growth via apoptosis pathway, however, the relation between them, the possible molecular mechanisms or the related downstream genes need further study.

Some studies have confirmed that LncRNAs maybe involved in the process of cancer cell proliferation, migration and invasion, et al [11, 17, 18]. However, the expression changes of LncRNA AC007392.4 have no effect on TSCC cells migration and invasion in the present study. Maybe LncRNA AC007392.4 is not involved in the migration and invasion process of TSCC cells.

In conclusion, our study revealed that the different expression of LncRNA AC007392.4 between TSCC and normal tissues and a possible role of LncRNA AC007392.4 in regulating the growth and apoptosis of TSCC cells, but not involved in the in the process of TSCC cells migration and invasion. Based on these findings, we suggest that LncRNA AC007392.4 may be an important and useful therapeutic biomarker for TSCC patients in future but need more study.

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Disclosure of conflict of interest

None.

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